POLYMERIC SCHIFF BASES. XVII. AZOMETHINE COPOLYMERS

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ABSTRACT

by solution processes. Melt polymerization techniques have been used to synthesize random, block and graft azomethine copolymers in which the repeating units correspond to units derived from either A-B monomers or a pair of A-A and B-B monomers. Not all systems yielded tractable copolymers, and these limitations are discussed. The block copolymers are prepared readily by the reaction of appropriate insoluble oligomers with monomers, or by the reaction of two different fusible oligomers. The thermal stabilities of the copolymers are high and comparable to azomethine homopolymers.

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INTRODUCTION

Various syntheses of monomeric and polymeric azomethines have been reported in a series of recent papers (1-20).

The direct condensation of aryldicarbonyl compounds with aryldiamines, or the self-condensation of aminoarylcarbonyl compounds, in solution and as melts, yielded, in most cases, low molecular weight, insoluble, infusible brick-dust polymeric azomethines. It was also shown (1,3-6,8,9) that propagation of these brick-dust products to higher molecular weight, tractable, black polymers could be accomplished in monomeric azomethines such as benzylideneaniline, which "dissolved" the products as a result of exchange reactions with the termini of the polymer chains:

OfHCAr'CH=NAr'N
$$\frac{1}{n}$$
H₂ $\frac{C_6H_5CH=NC_6H_5}{C_6H_5N_fHCAr'CH=NAr'N}$ HCC₆H₅ (eq. 1)

Alternately, black, tractable polymers could be synthesized by performing the condensation of the initial reagents in benzylideneaniline, or in the benzaldehyde and aniline components from which the benzylideneaniline was derived:

n OHCAr'CHO + n H₂NAr'NH₂
$$C_{6}H_{5}CH=NC_{6}H_{5}$$
 $C_{6}H_{5}CHO + C_{6}H_{5}NH_{2}$
 $C_{6}H_{5}N_{4}HCAr'CH=NAr'N_{5}HCC_{6}H_{5}$ (eq. 2)

The reactions of equation 2 were shown to be equivalent to the bis-exchange reaction:

n
$$C_{6}H_{5}N=HCAr'CH=NC_{6}H_{5} + n C_{6}H_{5}CH=NAr'N=HCC_{6}H_{5}$$

$$C_{6}H_{5}N=HCAr'CH=NAr'N=HCC_{6}H_{5} + (n-1) C_{6}H_{5}CH=NC_{6}H_{5} \qquad (eq. 3)$$

In all cases, propagation was the result of chain-end coupling with the regeneration of benzylideneaniline:

2
$$C_6H_5N_{+}HCAr'CH_NAr'N_{+}HCC_6H_5 \rightarrow$$

$$C_6H_5N_{+}HCAr'CH_NAr'N_{+}HCC_6H_5 + C_6H_5CH_NC_6H_5 \qquad (eq. 4)$$

Elimination of the benzylideneaniline was complicated by its retention apparently as a complex with the polymer in a solid polymer medium, and could be removed more completely at high temperatures at which it appeared, also, to coreact with the polymer (13-16).

The synthesis of high molecular weight polymeric azomethines would be simplified if the by-product of the condensation was a simple, small volatile molecule as water, such as would be formed by the interaction of $-\mathrm{NH}_2$ and $-\mathrm{CO}$ moieties located appropriately in the reactant molecules. Many attempts (1-3) to produce black polymeric asomethines by condensation reactions which eliminated water were not successful. However, it was observed that a number of high molecular weight, black polyketanils (10), $H_2 = NAr' C_{\frac{1}{2}} 0$, could be prepared in melt systems directly from the A-B type aminoaryl ketone monomers, HoNAr'COR, without the expedient of adding benzylideneaniline to the reaction mixture. Of the corresponding aminoarylaldehydes, HoNAr'CHO, only the orthoaminobenzaldehyde produced (10) homogeneous black polymeric melts, whereas infusible, intractable brick-dust trimers of melting points above 300°C were obtained from the condensations involving meta- and para-aminobenzaldehyde respectively. It was similarly noted that a number of black, high molecular weight polyketanils (8) could also be prepared directly in melt systems by the reaction of aryldiketones, RCOArCOR, with aryldiamines, H2NArNH2.

These observations led to this related study. First, consideration was given to the use of monomeric aminocarbonyl compounds as "solvents" for the readily prepared low molecular weight yellow-to-orange colored intractable, brick-dust polymeric azomethines. Since "solvents" of this type have reactive moieties which can react with the functional terminal groups of the brick-dust polymer chains without terminating the chains, chain propagation to higher molecular weight, tractable polymers could occur by the simultaneous elimination of water and block copolymer formation. Data on copolymeric azomethines,

whether block or random, appear to be absent in the literature. Accordingly, it was of a second interest to attempt the syntheses of tractable, random azomethine copolymers by using aminoarylcarbonyl monomers and aryl diketones as reactant "solvents", particularly in those polymerization systems whose reactants normally produce intractable, brick-dust polymeric azomethines. In these systems, "solvents" of this type would react statistically, without causing termination of the propagating chains, to yield random copolymers.

These studies were undertaken with the expectation that they would permit the syntheses of a wide variety of high molecular weight, block- and random-copolymeric azomethines which would pass through a fusible, or at least a tractable stage. The three block and two random copolymer systems investigated are classified as follows:

Type I-Block Copolymers

Equimolar amounts of a dicarbonyl compound, $Ar(CRO)_2$, and a diamino compound, $Ar(NH_2)_2$, can be considered as equivalent to the aminocarbonyl monomers, RH_2NArCOR, of equation 5, since, on reaction they yield H_2NArN=CArCOR intermediates, and so these systems were evaluated in this study, also.

Type II-Block Copolymers

This class of block copolymers is synthesized from pairs of fusible, tractable oligomers; some typical reactions can be generalized as:

As a modification of equation 10, similar to that mentioned for the type I-block copolymers, the fusible oligomer, $H_2 = NArC = 0$, can be reacted with equimolar portions of $Ar(COR)_2$ and $Ar(NH_2)_2$ instead of with the fusible A-A plus B-B type oligomer.

Type III-Block Copolymers

The synthesis of this class would involve reacting an infusible, high molecular weight, black polymeric azine with aminoarylcarbonyl monomers, which normally homopolymerized to tractable, fusible polymers, to determine whether or not they would become grafted to the infusible, polymeric azines. Conceivably, in such cases, grafting would occur by the reaction of the terminal -NH₂ or C=0 groups in the polymers with the complimentary H₂N- or -C=0 group in the H₂NArCOR monomers, or in their equivalent, equal molar quantities of Ar(COR)₂ with Ar(NH₂)₂. Accordingly, such copolymers are classified as Graft-Block Copolymers.

Graft copolymerization of this type would permit the fabrication of infusible polyazomethines by the simple expedient of dispersing the infusible polymer in a monomer or tractable polymer, which would act as a temporary plasticizer, and, then, forming the mass under heat and pressure to achieve graft-copolymerization simultaneously with the formation of a structure of the desired shape.

Random Copolymers

The polymers of this system would contain two different types of repeating units, and would be prepared directly from unpolymerized monomers. The simplest polymer would be one derived from two A-B type monomers (Class I) as given in equation 11.

$$n H_2NArCOR + n' H_2NAr'COR \rightarrow H_2\{NArC\} \frac{R}{n}\{NAr'C\} \frac{R}{n}$$
 (eq. 11)

Random copolymer should also result from the reaction of a multiplicity of A-A with B-B type monomers (Class II) such as:

2n RCOArCOR + n H₂NArNH₂ + n H₂NAr'NH₂ →

and

2n H₂NArNH₂ + n RCOArCOR + n RCOAr COR ---

as well as by a combination of A-B monomers with A-A plus A-B monomers, thus,

$$\begin{array}{ccc}
R & R & R \\
H_2 = NArC & \frac{1}{n} & NArN = CArC & \frac{1}{n} & O
\end{array}$$
(eq. 14)

In equations 11-14, the segmers represented by n and n' are located statistically in the chain and are shown in brackets only as a matter of convenience.

The copolymers of this study were prepared by melt procedures previously published (1,4,6,8-10). To contrast with the melt procedures, the syntheses of a number of copolymers were attempted using solution methods to determine whether or not the disorder, resulting in the polymer from the use of a plurality of monomers, would contribute some measure of solubility.

EXPERIMENTAL

1. Reactants.

a. Monomers. The purification or synthesis of the aryldiamines, aryldicarbonyl and the aminoarylcarbonyl compounds used in this study have been reported previously (2,8,10).

<u>b. Brick-dust Polyazomethines</u>. The synthesis of the intractable, brick-dust oligomeric azomethines used as reactants in this study have been reported previously or prepared by procedures previously published (1,8-11) and are given in Table 1.

Table 1

Data on Infusible Oligomeric Azomethines Used as Reactants

Oligomer Designa- tion	Structure	Color	Value* of n	Refer- ence
I-Y	$0 = HC - CH = N - CH = H_2$	yellow	4.0	1
II-Y	0-14 CH=N CH=N CH=N 1	yellow	4.8	1
IIİ-A	C6H2CHEN ON HCO CHIO	orange	5.1	1
IV-Y	$c_{6}H_{5}N_{+}HC_{+}O_{+}CH_{-}N_{-}O_{-}N_{+}^{-}H_{2}$	yellow	3.2	1
V-Y	c_{H_3} c_{H_3} c_{H_3} c_{H_2}	yellow	3.4	8
VI-Y	н ₂ ‡-⊙-сн <u>1</u> -по	yellow	3.0	10
VII-Y	H ₂ ^{£N} CH nO	yellow- brown	6.0	10
VIII-Y	$ \begin{array}{ccc} CH_3 & CH_3 \\ C-C & C-N-N & \frac{1}{n} & H_2 \end{array} $	yellow	2.0	11

^{*}Value estimated from analytical data.

c. Fusible, Low Molecular Weight Polyazomethines. Melt condensation procedures, previously published (1,8,10), were used to synthesize the fusible, tractable low molecular weight polymers used in this study. The condensations

were performed to the stage where the softening point of the polymers was at least 100°C, and the data are given in Table 2.

Table 2

Data on Fusible Oligomeric Azomethines Used as Reactants

Oligomer Designa- tion	Structure	Color	Softening Temperature °C	Refer- ence
I-F	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	brown	111	8
II-F	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	brown	133	8
III-F	H ₂ ^{£N} O C InO	brown	98	10
IV-F	$H_2 \neq N - \bigcirc - C \xrightarrow{CH_3} N$	dark brown	107	10
V-F	^C 6 ^H 5 H ₂ 4N-⊙-C-1 n 0	black	110	10
VI-F	н ₂ ±м-€	black	158	10

d. Infusible, Black Polyazomethines. Previously published melt-polymerization procedures (1,8,10) were used to prepare the insoluble, infusible high molecular weight polymers used in this study, and all of them were substantially insoluble in concentrated sulfuric acid at room temperature. The maximum temperature used in synthesis of the polymer, is designated according to previous codings (1) by the suffix H followed by the maximum temperature used. The data on the polymers used are given in Table 3.

Table 3

Data on Infusible Black Polyazomethines Used as Reactants

Polymer Designa- tion	Maximum Tempera- ture °C	Structure*	Refer- ence
I-B	н260	$0 \neq HC \bigcirc -CH = N \bigcirc -N \neq \frac{1}{n}H_2$	1
II-B	н400	$0 \neq HC \bigcirc -CH = N \bigcirc -N \neq \frac{1}{n}H_2$	1
III-B	н600	$0 \neq HC \bigcirc -CH = N \bigcirc -N \neq \frac{1}{n}H_2$	1
IV-B	н1176	O±HC·O-CH=N O-N-InH2	1
V-B	н380	c_{6} H ₅ CH $\frac{1}{n}$ O-N=HC- \bigcirc -CH $\frac{1}{n}$ O	1
VI-B	н400	$C_6H_5N_{HC} - CH = N - CH = M_nH_2$	1
VII-B	н400	$ \begin{array}{ccc} CH_3 & CH_3 \\ C + C - C - N - C - N + H_2 \end{array} $	8
VIII-B	н400	H ₂ =N-O-C=] _n O	10

^{*}These structures are idealized structures and the polymers which have been heated at 400°C or higher contain other structures resulting from inter- and intramolecular reactions (1,14-16).

2. Copolymerizations.

a. In Solution. The copolymerizations were performed using 0.02 mole as the total weight of monomer in 150 ml of solvent and 0.02 g of zinc chloride as a catalyst, when used. When benzene was used as the inactive solvent, the continuous azeotropic method was used (1). Dimethylacetamide (DMAC) was evaluated also, as an active solvent (1). Insoluble polymers were separated from DMAC by filtration; the soluble polymers were isolated by distillation of the DMAC from the reaction mass at 15 mm Hg pressure. Table 4 summarizes the data on the random copolymers prepared in solution.

b. Melt Copolymerizations. The melt procedure and the thermostatically controlled apparatus previously reported was used (1,4,6,8-10). An intimate mixture of the reactants was placed in the polymerization tube at room temperature and a nitrogen atmosphere established in the apparatus. The reaction tube

Table 4

Data on Random Copolymerizations in Solution

Poly-		Reactants		Mole	÷		%	Characteristic of
ner No.	А	В	C	Ratio A:B:C	Solvent"	Cata- lyst	Yield	Polymer
S-1	p-C ₆ H ₄ (CHO) ₂	m-С ₆ Н4 (СНО) ₂	m-C ₆ H ₄ (NH ₂) 2	1:1:2	В	No	001	yellow, brick-dust
S-2	m-C ₆ H ₄ (CHO) ₂	P-C ₆ H ₄ (CHO) ₂	$m-C_6H_4 ({ m NH}_2)_2$	1:1:2	В	No	86	yellow, brick-dust
S-3	р-С ₆ Н4 (СНО) 2	m-C ₆ H ₄ (NH ₂) 2	$p-c_6H_4 (NH_2)_2$	2:1:1	В	No	76	yellow, brick-dust
S-4	m-C ₆ H ₄ (CHO) ₂	m-C ₆ H ₄ (NH ₂) 2	p-C ₆ H ₄ (NH ₂) ₂	2:1:1	В	No	93	yellow, brick-dust
8-5	P-C ₆ H ₄ (CHO) ₂	P-C ₆ H ₄ (COCH ₃) ₂	m-C ₆ H ₄ (NH ₂) ₂	1:1:2	B	No	89	yellow, brick-dust
S-6	p-c ₆ H ₄ (CHO) ₂	p-C ₆ H ₄ (NH ₂) 2	o-H ₂ NC ₆ H ₄ CH0	1:1:1	æ	No	86	yellow, brick-dust
S-7	p-H2NC6H4CHO	o-H2NC6H4CHO	0 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	1:1:0	В	Yes	79	yellow, brick-dust
8-8	P-H2NC6H4COCH3	p-H ₂ NC ₆ H ₄ CHO		1:1:0	В	Yes	06	yellow, brick-dust
8-9	m-C ₆ H ₄ (CHO) ₂	p-c ₆ H ₄ (CHO) ₂	m-C ₆ H ₄ (NH ₂) ₂	1:1:2	DMAC	No	93	yellow, brick-dust
s-10	$^{\mathrm{m-C_6H_4}}(^{\mathrm{CHO}})_2$	p-C ₆ H ₄ (NH ₂) ₂	P-H ₂ NC ₆ H ₄ COCH ₃	1:1:1	DMAC	Yes	88	yellow, brick-dust
S-11	P-C ₆ H ₄ (COCH ₃) ₂	m-C ₆ H ₄ (NH ₂) ₂	p-C ₆ H ₄ (NH ₂) ₂	2:1:1	DMAC	Yes	91	yellow, brick-dust
s-12	P-H ₂ NC ₆ H ₄ COCH ₃	m-H ₂ NC ₆ H ₄ COCH ₃	# # # # # # # # # # # # # # # # # # #	1:1:0	DMAC	Yes	95	brown, soluble, fusible
s-13	p-C ₆ H ₄ (COCH ₃) ₂	m-C ₆ H ₄ (NH ₂) ₂	p-H ₂ NC ₆ H ₄ COCH ₃	1:1:1	DMAC	Yes	93	brown, soluble fusible

* B = benzene, DMAC = dimethylacetamide.

was then inserted in the aluminum heat-sink at 180°C and the reaction allowed to proceed according to the following schedule:

Temperature, °C	Hours
180	4
225	2
325	17
400	1

In a number of cases, samples were isolated during or at the end of the time periods at 225°C, 325°C and 400°C for evaluation, and they are designated in accordance with previous coding (1), as the H225, H325 and H400 polymers.

The polymerizations were attempted first as uncatalyzed reactions; when polymerization, as evidenced by an increase in melt viscosity, did not occur readily within one hour at 180°C, or was very sluggish, the reactions were repeated using 0.5 weight per cent of zinc chloride at the catalyst.

3. Thermogravimetric Analyses.

Thermogravimetric procedures previously published were used (1) and performed on a du Pont 900 Differential Thermal Analyzer in conjunction with a 950 Thermogravimetric Analyzer. The sample size was 10 mg ground to a powder of approximately 500 pieces; the rate of heating was 15°C per minute in a dry nitrogen or air stream at a gas flow-rate of one standard liter per minute to a recorded 1200°C temperature, which corresponds to 1176°C when corrected for the non-linearity of the chromel-alumel thermocouple.

4. Spectral Analyses.

A Perkin-Elmer 421 Grating Spectrophotometer was used to examine the samples prepared as potassium bromide discs.

5. Postheating of the Polymers.

A few samples of the copolymers were postheated to 600 and 1176°C by procedures previously published (1).

The formulations, elemental analyses and thermogravimetric analyses of the copolymers prepared are given in Tables 5 to 12 inclusive.

Table 5

Type I-Block Copolymers from Infusible Oligomers and A-B Type Monomers

					Analyses	vses Per	r Cent	
Polymer Designation	Oligomer	Monomer Used*	Zinc Chlorido	%*% V:012				H400
Eriachon	2000		מוזמי זמב	דבום	Ŋ	н	N	N
COP-1A	I-Y	o-H ₂ NC ₆ H ₄ COCH ₃	No	96	80.29	2.63	12.52	12.64
COP-2A	X-II	o-H2NC ₆ H4COCH3	No	86	80.31	5.58	12.49	12.68
COP-3A	X-III	o-H ₂ NC ₆ H ₄ COCH ₃	Yes	96	81.21	5.56	12.63	12.70
COP-4A.	IV-Y	$^{\mathrm{o-H}_2\mathrm{NC}_6\mathrm{H}_4\mathrm{COCH}_3}$	Yes	91	81.20	5.52	12.07	12.26
COP-5A	V-Y	$^{\mathrm{o-H}_{2}\mathrm{NC}_{6}\mathrm{H}_{4}\mathrm{COCH}_{3}}$	No	89	80.71	10.9	11.81	11.84
COP-6A	V-IV	o-H2NC ₆ H4COCH3	No	65	79.02	5.63	12.30	12.44
COP-7A	VII-Y	o-H ₂ NC ₆ H ₄ COCH ₃	No	65	79.10	5.66	12.34	12.43
COP-8A	VIII-Y	o-H ₂ NC ₆ H ₄ COCH ₃	No	93	76.56	6.35	14.31	14.52
COP-9A	I-Y	m-H ₂ NC ₆ H ₄ COCH ₃	Yes	100	80.43	5.52	12.50	12.67
COP-10A	Y-II	m-H ₂ NC ₆ H ₄ COCH ₃	No	66	80.36	5.61	12.53	12.69
COP-11A	I-Y	P-H2NC ₆ H4COCH3	No	86	80.33	5.53	12.48	12.59
COP-12A	I-Y	p-ch ₃ conhc ₆ h ₄ cho	Yes	96	80.80	4.98	13.43	13.49

* The molar amounts of monomer used corresponded to the value of n of the oligomer ** All reactions yielded clear, homogeneous melts or solutions at 180°C. so that feed was that of a 1:1 copolymer.

Table 6

Thermogravimetric Analyses in Nitrogen an Some Type I-Block Copolymers from Infusible Oligomers and Monomers

Polymer	Atmos-		<u> </u>	Per	Cent V	Veight	Loss	at °C	
Number	phere	400	500	600	700	800	900	1000	1176
COP-1A-H400 COP-1A-H600	nitrogen nitrogen	0.0	0.0	2.8 0.0	8.9 2.6	11.9 6.5	13.7 10.1	14.8 11.2	17.9 13.9
COP-2A-H400 COP-2A-H600	nitrogen nitrogen	0.0	0.0	2.4 0.0	4.6 1.8	8.0 6.2	13.5 8.7	14.5 10.8	17.6 13.2
COP-4A-H400 COP-4A-H600	nitrogen nitrogen	0.0 0.0	0.0	2.9 0.0	9.3 2.8	12.2 7.3	14.1 9.4	15.2 11.7	18.3 14.6
COP-6A-H400 COP-6A-H600	nitrogen nitrogen	0.0	0.0	3.2 0.0	9.5 3.0	12.5 7.2	14.5 9.3	15.6 11.4	18.8 16.2
COP-8A-H400 COP-8A-H600	nitrogen nitrogen	0.0	0.0	1.9	3.9 2.5	7.6 7.0	12.9 9.1	13.9 12.0	17.5 15.3
COP-9A-H400 COP-9A-H600	nitrogen nitrogen	0.0	0.0	3.0 0.0	9.6 2.6	12.7 6.2	14.8 8.2	15.7 11.5	18.7 14.9
COP-10A-H400 COP-10A-H600	nitrogen nitrogen	0.0	0.0	2.3 0.0	4.2 3.0	7.7 7.2	13.3	14.4 11.2	17.6 16.0
COP-11A-H400 COP-11A-H500	nitrogen nitrogen	0.0	0.0	2.5 0.0	4.8 1.9	8.2 6.5	13.8 8.9	14.8 10.9	17.8 14.3
COP-12A-H400 COP-12A-H600	nitrogen nitrogen	0.0	0.0	2.6 0.0	5.0 2.1	8.3 5.7	14.0 7.9	15.1 11.9	18.1 14.8

Table 7

Type I-Block Copolymers from Infusible Oligomers with A-A plus B-B Type Monomers

Polymer	01i-	Reactants	*	6	Observetions at		Analyses, %	es, %	
				° .	180°C	H	H325		H400
)	A-A Monomer	B-B Monomer	rieli	2 22	C	Н	N	N
COP-1B	X-IA	vi-т онс-О-сно	H ₂ N-(©)-NH ₂	65	clear initially, then brick-dust			-	
COP-2B	Y-IV	VI-Y OHC CHO	H ₂ N-CO NH ₂	97	partly in solution initially, then more heterogeneous, then brick-dust	-	-	8 8 8 8	# # #
COP-3B	X-IV	VI-Y H3COC CO-COCH3	H ₂ N-© NH ₂	76	clear solution and melt	81.19 5.76	5.76	12.42	12.44
COP-4B	VII-Y	VII-Y H3COC-O-COCH3	H ₂ N-O	92	clear solution and melt	80.97 5.72	5.72	12.39	12.46
COP-5B	VII-Y	VII-Y H3COC-O-COCH3	H2N-(O)-NH2	36	clear solution and melt	81.20	81.20 5.70	12.43	12.43

* The reactants were used in a 1:1 ratio in amounts which corresponded to the n value of the oligomer so that the feed would correspond to a 1:1 block copolymer.

Table 8

Thermogravimetric Analyses of Type I-Block Copolymers from Infusible Oligomers with A-A plus B-B Type Monomers

Polymer	Atmos-		Pei	Cent	Weight	Loss at	°C	
Number	phere	500	600	700	800	900	1000	1176
COP-3B-H400 COP-3B-H600	nitrogen nitrogen	0.0	2.6 0.0	4.9 0.0	8.2 1.4	13.8 6.6	14.7 10.3	17.8 14.0
COP-4B-H400 COP-4B-H600	nitrogen nitrogen	0.0	3.2 0.0	9.3 0.0	12.1	14.3 5.8	15.2 10.7	18.4 15.0
COP-5B-H400 COP-5B-H600	nitrogen nitrogen	0.0	3.0 0.0	9.0 0.0	12.0 2.0	13.9 7.7	14.9 10.3	18.1 15.4

Table 9

Block Copolymers from Mixtures of Fusible Oligomers

Copolymer	Reactan	ts*	Melt at		alyses	, %	
Designa-	Oligomer	Oligomer	180°C		Н325	· · · · · · · · · · · · · · · · · · ·	H400
tion	Oligomei	Oligomet		C	H	N	N
COP-1C	I-Y	VI-Y	No				
COP-2C	I-F	III-F	Clear	81.43	6.01	11.41	11.78
COP-3C	I-F	IV-F	Clear	81.37	6.00	11.70	11.84
COP-4C	I-F	V-F	Clear	81.41	5. 97	11.78	11.85
COP-5C	II-F	IV-F	Clear	80.81	5.51	12.59	12.67
COP-6C	II-Y	VI-F	Clear	83.64	5.53	9.82	9.93
COP-7C	I-Y	IV-F	No				
COP-8C	IIIF	IV-F	Clear	81.04	6.11	11.77	11.88
COP-9C	III-F	V-F	Clear	83.30	5.61	9.79	9.86
COP-10C	I-F	VIII-F	Clear	81.79	5.96	11.90	11.94

^{*}Reactants were used in amounts equivalent to a 1:1 copolymer.

Table 10

Thermogravimetric Analyses of Block Copolymers Prepared from Mixtures of Oligomer

Copolymer	Atmosphere		H	er Cent	Weight	Loss at	°C	
Designation	Acmosphere	500	600	700	800	900	1000	1176
COP-2C-H400	nitrogen air	0.0	2.7 69.3	5.1 100.0	8.4	13.7	14.7	16.9
COP-3C-H400	nitrogen air	0.0 3.4	2.1 71.8	4.5 100.0	9.8	13.8	15.1	17.1
СОР-4С-Н400	nitrogen air	0.0 3.7	3.1 72.6	5.7 100.0	11.6	14.4	15.2	18.3
COP-5C-H400	nitrogen air	0.0 3.3	2.4 69.7	4.7 100.0	8.1	13.6	14.7	17.5
СОР-6С-Н400	nitrogen air	0.0 2.7	2.3 71.3	4.8 100.0	10.1	14.2	15.8	17.4
COP-8C-H400 COP-8C-H600 COP-8C-H1176	nitrogen nitrogen nitrogen	0.0 0.0 0.0	2.6 0.0 0.0	8.3 1.9 0.0	11.6 5.8 0.0	13.5 7.8 0.0	14.4 10.7 0.0	17.6 14.8 2.0
COP-9C-H400 COP-9C-H600 COP-9C-H1176	nitrogen nitrogen nitrogen	0.0 0.0 0.0	2.9 0.0 0.0	8.0 3.3 0.0	9.8 7.5 0.0	14.8 9.5 0.0	15.9 11.1 0.0	21.1 16.0 3.3
COP-10C-H400	nitrogen air	0.0 3.6	2.9 69.9	8.2 100.0	9.7	14.0	15.1	18.5

Table 11

Type II-Block Copolymers from Fusible Oligomers with A-A Plus B-B Type Monomers

Polvmer	011-	Reactant	nts*	6	Observation	Elem	ental.	Elemental Analyses	
		A-A	B-B	V:61d	24 180°C		H325		005H
Number	gomer [Monomer	Monomer	34 34 4	מר זמת ת	၁	н	N	Z
COP-1D	III-F	ит-ғ онс-О-сно	$H_2N \bigcirc M_2$	16	heterogeneous, then brick-dust				8 8
COP-2D	III-F	111-ғ н3сос-©-сосн3	$^{\mathrm{NH}_2}$	96	clear solution and melt	81.26	6.04	81.26 6.04 11.85	11.90
COP-3D	V-F	V-F H3COC-O-COCH3	NH2 (○)-N2	76	clear solution and melt	81.31	10.9	81.31 6.01 11.87	11.93
COP-4D	VI-F	VI-F H3COC O-COCH3	H ₂ N $\langle \bigcirc \rangle$ NH ₂	93	clear solution and melt	82.99	5.70	82.99 5.70 10.51	10.58
COP-5D	VII-F	VII-F H3COC-©-COCH3	^{NH2} (○)- ^{NH2}	67	clear solution and melt	81.26	5.74	81.26 5.74 12.40	12,45

* A-A and B-B monomers were used in a 1:1 molar ratio in amounts with the oligomer to give a 1:1 copolymer.

Table 12

Thermogravimetric Analyses of Type II-Block Copolymers from Fusible Oligomers with A-A Plus B-B Type Monomers

Polymer	Atmosphere		Pe	r Cent	Per Cent Weight Loss at °C	Loss a	ς C	
Number	a saud comar	500	009	200	800	006	1000	1176
COP-2D-H400	nitrogen	0.0	3.2	3.2 9.6	12.4	14.5	15.7	18.8
COP-2D-H600	nitrogen	0.0	0.0 0.0	0.0	6.5	8.9	11.4	15.3
COP-3D-H400	nitrogen	0.0	2.9 9.7	7.6	12.7	14.4	15.9	18.0
COP-3D-H600	nitrogen	0.0	0.0	0.0	5.7	8.1	10.7	14.5
COP-4D-H400	nitrogen	0.0	2.1	4.4	6.7	13.4	14.4	17.4
COP-4D-H600	nitrogen	0.0	0.0	0.0	6.3	8.3	11.1	14.9
COP-5D-H400	nitrogen	0.0	1.1	3.8	7.5	13.0	14.2	16.3
COP-5D-H600	nitrogen	0.0	0.0	0.0	1.0	5.3	10.1	13.8

c. Graft-Block Copolymerizations. The general procedure consisted in grinding the infusible, black polyazomethines to a very fine powder, to which was added the monomer, and zinc chloride, if used. The mixture was then blended to uniformity, in a Spex-Mixer-Mill, introduced into the polymerization tube, a nitrogen atmosphere established in the tube which was then inserted into the metal heat-sink preheated to and controlled at 180°C. The polymerization schedule used was

Temperature, °C	Time:hours
180	1
225	1
300	2
400	8

Duplicate experiments were performed for a number of these copolymerizations so that samples could be withdrawn at the end of the 300°C heating period and extracted with dimethylacetamide and acetone. The infusible polymer and monomer were used in a ratio of 1 gm of polymer to 4 grams of monomer. None of the mixtures melted during the course of the reaction; at the end of the reaction, the product was isolated as a compact integral button. The data for these polymerizations are summarized in Tables 13 and 14.

Table 13

Data on Graft Copolymers from Infusible Polyazomethines and Aminoarylcarbonyl Monomers

Graft	Reactants	s Used	2	**%	% Yield	Ana 1 H400	Analyses of H400 Polymers.	%
Designation	Polymer*	Monomer	4nc12	at 300°C	at 400°C	ပ	н	1 1
GC-1	I-B-H260	m-H2NC6H4COCH3	No	205	363	81.01	5.98	12.28
GC-2	II-B-H400	m-H ₂ NC ₆ H ₄ COCH ₃	Yes	238	429	80.93	10.9	12.06
60-3	III-B-H600	m-H2NC6H4COCH3	yes.	262	4,08	81.21	6.10	12.43
7-D5	IV-B-H1176	176 m-H ₂ NC ₆ H ₄ COCH ₃	Yes	248	437	89.08	5.88	12.19
GC~5	V-B-H380	o-H ₂ NC ₆ H ₄ CHO	Yes	226	438	80.66	5.25	12.98
925	VI-B-H400	o-H2NC6H4CHO	Yes	243	421	81.11	6.07	12.01
GC-7	VII-B-H400	p-H2NC6H4COCH3	Yes	222	427	81.10	6.33	11.91
8-25	VIII-B-H400	o-H ₂ NC ₆ H ₄ CHO	No	197	378	80.68	6.07	12.23
6-25	VII-B-H400	o-H2NC ₆ H4COCH3	Yes	228	414	81.09	6.28	12.01
GC-10	VII-B-H400	o-H ₂ NC ₆ H ₄ COC ₆ H ₅	Yes	239	432	99• 78	5.27	8.82

**Based on weight of infusible polymer* used as 100%.

Table 14

Thermogravimetric Analyses of Some Graft H400-Copolymers

Copolymer	Atmosphere		1	er Cent	: Weight :	Loss at	°C		
Designation	Acmosphere	400	500	600	700	800	900	1000	1176
GC-1-H400	nitrogen air	0.0	0.0 2.6	2.7 70.1	8.8 100.0	12.0	13.9	15.0	18.1
GC-3-H400	nitrogen air	0.0	0.0 3.5	3.0 72.8	9.2 100.0	12.1	14.2	15.2	18.4
GC-4-H400	nitrogen air	0.0	0.0 3.9	2.8 72.7	4.7 100.0	8.1	13.5	15.4	18.8
GC-5-H400	nitrogen air	0.0	0.0 3.0	2.3 71.7	4.7 100.0	7.9	13.6	14.6	17.7
GC-6-H400	nitrogen air	0.0	0.0 3.3	2.4 71.0	4.9 100.0	8.3	13.9	14.9	18.0
GC-7-H400	nitrogen air	0.0	0.0 2.9	2.9 68.5	8.6 100.0	11.6	14.1	15.0	18.3
GC-8-H400	nitrogen air	0.0	0.0 3.1	2.6 69.1	5.0 100.0	8.3	1.3.9	14.8	17.9
GC-10-H400	nitrogen air	0.0	0.0 2.4	1.4 70.2	4.1 100.0	7.9	13.3	14.5	16.9

DISCUSSION

The reactants for the syntheses of the copolymers of this study were selected so that in all cases, the by-product of condensation would be a simple molecule such as water, the only exception being the use of formyl acetanilide which eliminated acetic acid readily. Preliminary studies indicated that they could be prepared relatively easier by the exchange reactions, particularly by the bis-Schiff base exchange reactions (1), but, this method was not used primarily to avoid the complications of inter- and intramolecular reactions that occur in polymer systems containing benzylideneaniline (1,9,14-16).

The statistically large number of copolymer compositions possible from the large number of monomers available (1,6,8,10) for this study required first that a choice be made of the monomers to be used. Accordingly, the relatively simple, available monomeric reactants shown in Table 4 were selected. Even with a small

number of monomers, the possible number of copolymers is large since the number of copolymer compositions, from reactants which can produce two different repeating units, is very high between the range of 100 molar per cent of one repeating unit to 100 molar per cent of the second repeating unit. Thus, it was decided that those copolymers which contained only two types of repeating units in a 1:1 molar ratio would permit ready comparison between the various copolymers; so the monomer feed corresponding to this 1:1 composition was used in the syntheses. Copolymerizations in Solutions.

Copolymerizations in solution were undertaken first using a relatively inactive solvent such as benzene, and an active solvent such as dimethylacetamide (DMAC), to determine if soluble or fusible copolymers could be thus prepared. These copolymers are designated as the S-series shown in Table 4.

As a rule, the random copolymerizations in solution of the A-A plus B-B monomers yielded intractable brick-dust polymers whether the solvent was benzene or DMAC, or whether a pair of isomeric A-A aldehyde monomers were reacted with two moles of B-B diamine monomer, or a pair of B-B diamine monomers were reacted with two moles of A-A aldehyde monomers as evident in polymers S-1 to-S-4 and S-9. In general, it appeared that if the individual pairs of the A-A plus B-B reactants yielded intractable, brick-dust polymers, the copolymers were obtained also as brick-dusts since the disorder in the chain was insufficient to contribute solubility even in DMAC. Similarly, brick-dust copolymers were obtained when even greater linear disorder was introduced in the polymer chain by substituting half the molar quantity of A-A dialdehyde monomer by an A-A diketone monomer in the reaction with a B-B diamine as in polymer S-5. Similarly, complete substitution of the dialdehyde in Polymer S-3 by a diketone in the reaction with a pair of diamines as in Polymer S-11 also yielded an intractable brick-dust polymer. Modification of the A-A dialdehyde plus B-B diamine reaction by an aminoaryl carbonyl monomer as in polymers S-6 and S-10 also failed to yield tractable polymers. Also the reaction pairs of aminoaldehydes in benzene, as in S-7, or the reaction of an aminoaldehyde with an aminoketone in benzene, as in S-8, yielded intractable polymers. In contrast, the copolymerizations in DMAC of two aminoketones in S-12, or the coreaction of an A-A diketone plus a B-B diamine with an aminoketone as in S-13, yielded soluble, fusible polymers; these results were not unexpected, since (a) the individual low molecular weight polymers of the para- and meta-aminoacetophenones respectively are very soluble in DMAC (10), and (b) the low molecular weight polymers from p-diacetylbenzene and m-phenylenediamine are slightly soluble in DMAC (8), which solubility is increased by copolymerization with p-aminoacetophenone. In general, it may be concluded that copolymerization in solution does not yield tractable polymers if the monomeric components normally produce individual intractable homopolymers in that specific solvent.

Preliminary experiments, in solution of such reactant pairs as (a) infusible oligomers (Table 1) with A-B type monomers and (b) pairs of fusible oligomers (Table 2) also yielded intractable products. In view of the broad inapplicability of solution polymerizations for the syntheses of polyazomethines, the melt process was used to attempt the syntheses of the copolymers.

Melt Copolymerizations.

Type I-Block Copolymers. From Infusible Oligomers and A-B Type Monomers.

These were prepared by the melt process and are designated as the COP-A Series in Table 5. The feed composition for these copolymers, shown in Table 5, corresponds to that required for a 1:1 molar ratio of two different repeating units and thus offers a comparison with the S-series of copolymers made in solution. The reaction mixtures in the COP-series yielded clear, homogeneous melts by the end of the heating period at 180°C, and at the end of the 225°C heating period, the intrinsic viscosities in 98% H₂SO₄ were in the range of 0.09 to 0.11 d1/g and the softening points of the low molecular/copolymers ranged from 190 to 245°C. The initial intrinsic viscosities of the oligomers used were in the

range of 0.04 to 0.55 (1), and under similar thermal treatment, in the absence of zinc chloride, the A-B monomers used do not polymerize, while in the presence of zinc chloride, under the same conditions, the intrinsic viscosities in ${\rm H_2SO_4}$ of the homopolymers from the A-B monomers range from 0.02 to 0.04 d1/g. The solubilization of the oligomers and the intrinsic viscosity data indicate that copolymerization had occurred. Since the A-B monomers and their homopolymers formed at 225°C are soluble in acetone (10), whereas the oligomers used are insoluble in acetone (1), samples of the H225 polymers were extracted with acetone and dried at 225°C and the increase in weight as a function of the original weight of the oligomer was in the range of 7 to 16%. As the polymerization proceeded, the melt viscosity of the copolymer increased while the color changed progressively from yellow to brown to dark brown while water of condensation was eliminated. All the copolymers were hard, black glassy polymers at the end of the 325°C heating period and were insoluble in concentrated sulfuric acid. Samples withdrawn from the reactions of polymers COP-1A and COP-5A after two hours of heating at 325°C had intrinsic viscosities in concentrated sulfuric acid of 0.32 and 0.27 d1/g respectively. The elemental analyses of the H325 polymers are in good agreement with the calculated values for the 1:1 copolymers, which would indicate that the polymers had not undergone inter- and intramolecular reactions to any great extent, as occurs readily when the polymers are prepared by exchange reactions (1) particularly those which eliminated an amine or a Schiff base (15,16). The infrared spectrum of the H325 showed the presence of bands for -C=N- in the region 1600 cm⁻¹ with some evidence for the C=O function in the 1700 ${\rm cm}^{-1}$ region and ${\rm NH}_2$ in the 3380 ${\rm cm}^{-1}$ region; the spectra for the H400 polymers showed the bands for the C=N linkages but not for the NH $_2$ and C=O functions. The spectra of these copolymers corresponded to those obtained by superimposing the spectra (1,8,10) of the oligomers used on the spectrum of the polymerized A-B monomer, for example, the spectrum of COP-1A appeared to be the

The thermal stabilities of representative H400 and H600 Type I-Block Copolymers are given in Table 6 and, in general, are comparable to the stabilities of typical H400 and H600 polyazomethines previously reported (1,5-9,12-16).

Type I-Block Copolymers. From Infusible Oligomers with A-A plus B-B Type Monomers.

These also were prepared by the melt process and the pertinent data are given in Tables 7 and 8. Homogeneous melts were obtained at 180°C only in those systems in which the A-A plus B-B monomer pairs themselves would homopolymerize to tractable polymers as evidenced by copolymers number COP-3B-COP-5B inclusive. This was in contrast with systems COP-1B and COP-2B which showed some measure of initial homogeneity before becoming heterogeneous and yielded brick-dust polymers. The brick-dust polymers were not characterized further. The analytical values of the copolymers which passed through a clear melt stage are in good agreement with the calculated values, and the thermal properties given in Table 8 are typical values for typical H400 and H600 polyazomethines previously reported (1,5-9,12-16).

Type II-Block Copolymers. From Pairs of Fusible Oligomers.

This class of copolymers was synthesized from pairs of fusible oligomers of the type shown in Table 2, and the pertinent data on these copolymers are given in Tables 9 and 10. These copolymers are designated as the COP-C series. Included, for contrast, in the studies of Table 9 are attempts to prepare COP-1C from a pair of infusible oligomers, and COP-7C from a mixture of a fusible and an infusible oligomer; both efforts were unsuccessful. The syntheses of block copolymers from fusible oligomers were more facile, as would be expected, than the synthesis of the Type I-Block Copolymers from an infusible oligomer with an A-B type or with A-A plus B-B type monomers. The elemental analyses of the block copolymers

are in good agreement with the calculated values for the 1:1 copolymers, and their infrared spectra corresponded as in the Type I-Block Copolymers to the super-imposition of the spectra of the homopolymers of the two reacting components. The thermal stabilities of this class of block copolymers are summarized in Table 10 and they are comparable to the stabilities of typical polymeric azomethines (1,5-9,12-16).

Type II-Block Copolymers. From Fusible Oligomers with A-A Plus B-B Monomers.

The pertinent data are given in Tables 11 and 12, and these copolymers are designated as the COP-D series. As in the case of the Type I-Block Copolymer series, only those systems were successful in which the A-A with B-B monomers normally would yield fusible homopolymers as seen for COP-3D to COP-5D inclusive. The system for COP-1D was not successful, though it had been anticipated that a homogeneous melt would be obtained due to the solubilizing effect of the fusible oligomer III-F. Such was not the case, and the result confirmed the similar results obtained for the Type I-Block Copolymers, COP-1B and COP-2B. The elemental analyses shown in Table 11 of the copolymers which yielded homogeneous melts and their thermal analytical data, shown in Table 12, are typical of polymeric azomethines.

Type III-Block Copolymers. From Infusible Polymers with A-B Monomers.

The systems evaluated for the synthesis of this class of copolymers and the pertinent data are given in Tables 13 and 14. The per cent yields given in Table 13 have been calculated on the amount of initial infusible polymer as the 100% basis. The yield values of the H300 polymers at 300°C were obtained by extracting samples of the polymers first with DMAC, then with acetone followed by drying at 200°C. These extractions removed unreacted monomers and low molecular weight homopolymers, if any, of the aminoketones, which may have formed. The theoretical yield for the H400 polymers is in the range of 445-447% on the basis of initial infusible polymer. The yields of the uncatalyzed reactions were lower

than those which were catalyzed by ZnCl₂, and the lower than theoretical yield in all cases is attributable to sublimation losses of the monomer. The thermogravimetric data of Table 14 is representative of typical polymeric azomethines. The interesting observation made in these studies is that integral castings were obtained, in most cases, at a ratio of 4 parts of monomer to 1 part of polymer, and not at lower ratios of monomer to polymer. At the lower ratios the monomer appeared to be absorbed by the polymer and to become unavailable to function as an adhesive.

Type I-Random Copolymers.

The unsuitability of the solution method for preparing random copolymers led to the use of melt polymerizations in attempts to prepare these copolymers according to equations 11 to 14 inclusive. The random copolymerization of the type represented by equation 11 was most facile when the A-B monomers were the aminoaryl ketones, each of which homopolymerized to tractable polymers, or when one A-B monomer was an aminoketone and the other A-B monomer was an aminoaldehyde. However, an intractable brick-dust polymer was obtained when both A-B monomers were aldehydes. These copolymers are designated as RC-A series and the pertinent data are given in Tables 15 and 16.

The random copolymerizations of the type represented by equations 12 and 13 and designated as the RC-B series, were also very facile and tractable melts were obtained when the dicarbonyl compounds, RCOArCOR were the diketones, as shown by the copolymers RC-9B to RC-12B inclusive in Table 17. In contrast, only brick-dust polymers were obtained when the dicarbonyl compounds were dialdehydes, as represented by copolymers RC-3B, RC-4B and RC-5B, in which a 2.5:7.5 ratio of repeating units were introduced instead of the 1:1 ratio. When an equimolar mixture of a dialdehyde, Ar(CHO)₂ and a diketone, Ar(COCH₃)₂ as represented by copolymer RC-1A, only brick-dust polymers were obtained and tractable systems were obtained only when the ratio of diketone to dialdehyde was 7.5 to 2.5 as in copolymers TC-6B, RC-7B and RC-2B, and in the case of RC-2B, the product was a granular, brick-dust intractable polymer at the end of the 225°C heating period. The elemental and thermogravimetric analyses of polymers RC-6B to RC-12B, which in the course of synthesis, remained homogeneous, are given in Tables 18 and 19.

Table 15

Random Copolymers from Pairs of A-B Monomers

Polymer		Reactants*	ants*		Observation		Analyses, %	es, %	
Nimber					at 180°C		H325	5	H400
Namper	Monomer	Mole	Monomer	Моте		ט	ਸ਼	N	Z
RC-1A	^{Н2N} ⊙-сосн ₃	1.0	н ₂ и-(⊙}-сосн ₃	1.0	melt	80.83	6.07	11.83	11.93
RC-2A	NH2 ©>-cocH3	1.0	H2N O COCH3	1.0	melt	80.73	6.11 11.81	11.81	11,91
RC-3A	NH2 (©)-cocH ₃	1.0	н ₂ м-⊙-сосн ₃	1.0	melt	80.91	5.99	11.77	11.89
RC-4A	н2 ^N С>сосн3	1.0	H ₂ N-COC ₆ H ₅	1.0	melt	83.51	5.66	9.82	9.88
RC-5A	O-CHO	1.0	O COCH ₃	1.0	melt	80.57	5.58	12.60	12.68
RC-6A	⊙ CHO	1.0	H ₂ N-(©)-coch ₃	1.0	melt	81.01	5.60	12.59	12.66
RC-7A	NH2 CHO CHO	1.0	H ₂ N (©)-CHO	1.0	brick-dust	1 1 1 1	! ! !	1 1 1 1	: : :

* All reactions were catalyzed by zinc chloride.

Table 16

Thermogravimetric Analyses of Some Random Copolymers from Pairs of A-B Monomers

Polymer	Atmos-		1	er Cen	t Weigh	t Loss a	t °C	
Number	phere	500	600	700	800	900	1000	1176
RC-1A-H400	nitrogen	0.0	3.0	8.3	9.8	14.1	15.2	18.3
-H600	nitrogen	0.0	0.0	1.9	6.4	8.8	10.8	13.8
-H1176	nitrogen	0.0	0.0	0.0	0.0	0.0	0.0	1.8
RC-2A-H400	nitrogen	0.0	2.4	4.6	9.7	13.7	14.9	18.4
-H600	nitrogen	0.0	0.0	2.1	5.6	7.8	11.7	14.9
-H1176	nitrogen	0.0	0.0	0.0	0.0	0.0	0.0	2.1
RC-3A-H400	nitrogen	0.0	2.7	5.2	9.4	14.3	15.6	17.4
-H600	nitrogen	0.0	0.0	3.0	7.3	9.3	11.3	15.9
-H1176	nitrogen	0.0	0.0	0.0	0.0	0.0	0.0	2.4
RC-5A-H400	nitrogen	0.0	2.6	8.4	11.7	13.7	14.8	17.9
-H600	nitrogen	0.0	0.0	2.6	5.9	8.5	11.1	15.0
-H1176	nitrogen	0.0	0.0	0.0	0.0	0.0	0.0	2.8

Random Copolymers from a Pair of A-A Type Monomers with a Single B-B Type Monomer Table 17

Polymer	Reactant	Mole	Reactant	Mole	Reactant	Mo1e		Observations at '	0 °
Number							180	225	325
RC-1B	онс-О-сно	0.50	н3сос-⊙-сосн3	0.50	$H_2N \bigcirc NH_2$	1.0	melt	brick-dust	
RC-2B	оно-{О}-ноо	0.25	н3сос-О}-сосн3	0.75	H ₂ N-{O}-NH ₂	1.0	melt	brick-dust	
RC-3B	осн-О-сно	0.50	онс-О-сно	0.50	H ₂ N-∰-NH ₂	1.0	melt	brick-dust	1 1 1 1 1 1 1
RC-4B	онс-О-сно	0.50	оно-Оно	0.50	H ₂ N O-NH ₂	1.0	melt	brick-dust	
RC-5B	онс-О-сно	0.25	ОНС ОНС	0.75	H ₂ N CO-NH ₂	1.0	melt	brick-dust	
RC-6B	онс-СО	0.25	н3сос-⊘-сосн3	0.75	H ₂ N-(©)-NH ₂	1.0	melt	very viscous	black glassy
RC-7B	оно Со	0.25	н3сос-⊙-сосн3	0.75	H2N O-NH2	1.0	melt	very viscous	black glassy
RC-8B	онс-СУ-сно	0.25	нзсос-О-соснз	0.75	$^{\mathrm{H}_{2}\mathrm{N}}\left\langle \bigcirc\right\rangle ^{\mathrm{NH}_{2}}$	1.0	melt	very viscous	black glassy
RC-9B	н3сос-О-сосн3	0.50	(H3COC (O)-)-20	0.50	H ₂ N-(O)-NH ₂	1.0	melt	very viscous	black glassy
RC-10B	$(H_3ccc \cdot \bigcirc) - \frac{1}{2}$	0.50	о ∠(-{⊙}-)	0.50	H ₂ N-(C)-NH ₂	1.0	melt	very viscous	black glassy
RC-11B	(H3coc-{○}- }_2 cH2	0.50	$(H_3\cos{\langle \bigcirc \rangle})_{\overline{\Sigma}}$ 80 $_2$.50	H ₂ N-(©)-NH ₂	1.0°	melt	very viscous	black glassy
RC-12B	(H3COC-(O)-)2CH2 0.50	0.50	н3сос-(⊙)-сосн3	0.50	H2N O-NH2	1.0	melt	very viscous	black glassy

Table 18

Elemental Analyses of the Tractable RC-B Series of Random Copolymers

Polymer	A	nalysi	s, %	
Number		Н325	-	H400
Treations 2	С	H	N	N
RC-6B	80.88	5.37	12.66	12.71
RC-7B	80.79	5.42	12.63	12.69
RC-8B	80.81	5.51	12.70	12.74
RC-9B	80.77	5.70	10.16	10.20
RC-10B	82.24	5.82	8.52	8.54
RC-11B	77.19	5.33	7.99	8.02
RC-12B	82.94	5.99	10.22	10.27

Table 19

Thermogravimetric Analyses of Some of the RC-B Series of Random Copolymers in Nitrogen and in Air

Polymer	Atmos-		P	er Cent	Weight	Loss a	t °C		
Number	phere	400	500	600	700	008	900	1000	1176
RC-6B-H400 RC-6B-H400	nitrogen air	0.0	0.0 6.2	2.5 100.0	4.7	8.0	13.6	14.5	17.6
RC-7B-H400 RC-7B-H400	nitrogen air	0.0	0.0 6.3	2.9 100.0	9.0	12.0	13.8	14.9	18.0
RC-8B-H400 RC-8B-H400	nitrogen air	0.0	0.0 7.4	3.0 100.0	9.3	12.2	14.2	15.2	18.4
RC-9B-H400 RC-9B-H400	nitrogen air	0.0	0.0 6.8	3.3 100.0	9.6	12.5	14.6	15.6	18.9
RC-10B-H400 RC-10B-H400	nitrogen air	0.0	0.0 8.1	3.1 100.0	9.7	12.7	14.8	15.8	18.8
RC-11B-H400 RC-11B-H400	nitrogen air	0.0	0.0 7.7	1.9 100.0	4.0	7.7	13.0	14.0	17.4
RC-12B-H400 RC-12B-H400	nitrogen air	0.0	0.0 7.3	2.3 100.0	4.2	7.8	13.4	14.3	17.5

The random copolymerizations represented by equation 14 were also very facile and homogeneous, tractable melts and were obtained when the dicarbonyl compound was a diketone, CH₃COArCOCH₃, in which case the aminocarbonyl compound, H₂NArCOR, could be either a ketone or an aldehyde; such products are represented by copolymers RC-8C to RC-11C inclusive of Table 20.

When the dicarbonyl compound in equation 14 was a dialdehyde, C_6H_4 (CHO)₂, intractable condensation products were obtained unless four or more moles of aminoketone were used, as shown by copolymer numbers RC-1C to RC-7C inclusive. However, even when four moles of an aminoaldehyde were used instead of the aminoketone, as shown by copolymer RC-12C, a brick-dust product was obtained.

Elemental analyses and the thermogravimetric data of the RC-C Series of copolymers are given in Tables 21 and 22 and are typical of the polymeric azomethines.

Random Copolymers from Mixtures of A-A, B-B and A-B Type Monomers Table 20

Number Monomer Mole RC-1C OHC-⊙-CHO 1.0 H2 RC-2C OHC-⊙-CHO 1.0 H2 RC-3C OHC-⊙-CHO 1.0 H2 RC-4C OHC-⊙-CHO 1.0 H2 RC-5C OHC-⊙-CHO 1.0 H2 RC-5C OHC-⊙-CHO 1.0 H2 RC-6C OHC-⊙-CHO 1.0 H2 RC-7C OHC-⊙-CHO 1.0 H2 RC-8C H3COC-⊙-CHO 1.0 H2 RC-9C H3COC-⊙-COCH3 1.0 H2 RC-10C H3COC-⊙-COCH3 1.0 H2 RC-11C H3COC-⊙-COCH3 1.0 H2		: נ				277073377000) 1
OHC-←○-CHO 1.0 H ₃ COC-←○-COCH ₃ 1.0	e Monomer	Mole	Monomer	Mole	180	225	325
OHC-♠-CHO 1.0 OHC-♠-CHO 1.0 OHC-♠-CHO 1.0 OHC-♠-CHO 1.0 OHC-♠-CHO 1.0 H3COC-♠-COCH3 1.0 H3COC-♠-COCH3 1.0 H3COC-♠-COCH3 1.0	$0 \mid H_2^{N-\bigcirc NH_2}$	1.0	н ₂ м-⊖-сосн ₃	3.0	paste	granular	partly glassy
OHC-♠-CHO 1.0 OHC-♠-CHO 1.0 OHC-♠-CHO 1.0 OHC-♠-CHO 1.0 H3COC-♠-COCH3 1.0 H3COC-♠-COCH3 1.0 H3COC-♠-COCH3 1.0	$0 \mid H_2^{N} \bigcirc M_2$	1.0	H ₂ N-(©)-cocH ₃	3.0	melt	granular	black granular
OHC-(○)-CHO 1.0 OHC-(○)-CHO 1.0 OHC-(○)-CHO 1.0 H ₃ COC-(○)-COCH ₃ 1.0 H ₃ COC-(○)-COCH ₃ 1.0 H ₃ COC-(○)-COCH ₃ 1.0	$0 \mid H_2N+\bigcirc -NH_2$	1.0	н2м-⊘-сосн3	0*5	melt	viscous	black glassy
OHC-O-CHO 1.0 OHC-O-CHO 1.0 OHC-O-CHO 1.0 H ₃ COC-O-COCH ₃ 1.0 H ₃ COC-O-COCH ₃ 1.0 H ₃ COC-O-COCH ₃ 1.0	0 H ₂ N-©-NH ₂	1.0	H ₂ N ⊕ COCH ₃	4.0	melt	viscous	black glassy
OHC-☉-CHO 1.0 OHC-☉-CHO 1.0 H ₃ COC-☉-COCH ₃ 1.0	$0 \mid H_2^{N} + \bigcirc M_2$	1.0	H ₂ N-€)-cocH ₃	4.0	melt	viscous	black glassy
OHC T⊕ CHO 1.0 H ₃ COC ← C COCH ₃ 1.0 H ₃ COC ← C COCH ₃ 1.0 H ₃ COC ← C COCH ₃ 1.0	0 H2N-O-NH2	1.0	H ₂ N-COCH ₃	4.0	melt	viscous	black glassy
H ₃ COC-⊙-COCH ₃ 1.0 H ₃ COC-⊙-COCH ₃ 1.0 H ₃ COC-⊙-COCH ₃ 1.0	$0 \mid ^{\mathrm{H}_{2}\mathrm{N}} \bigcirc ^{\mathrm{NH}_{2}}$	1.0	н ₂ м-О-сос ₆ н ₅	4.0	melt	viscous	black glassy
H ₃ COC (○) COCH ₃ 1.0 H ₃ COC (○) COCH ₃ 1.0 H ₃ COC (○) COCH ₃ 1.0	$0 \mid H_2^{N-\bigcirc NH_2}$	1.0	н₂м-⊘-сосн₃	2.0	melt	viscous	black glassy
H ₃ COC-(○)-COCH ₃ 1.0 H ₃ COC-(○)-COCH ₃ 1.0	$0 \mid H_2^{\text{N}} \bigodot^{\text{NH}_2}$	1.0	H ₂ N-(C)-cocH ₃	1.0	melt	viscous	black glassy
н₃сос-⊙-сосн₃ 1.0	$0 \mid H_2^{N} \oplus M_2$	1.0	H ₂ N-(C)-coc ₆ H ₅	1.0	me1t	viscous	black glassy
	O H2N-©-NH2	1.0	ин ₂ О)-сно	1.0	melt	viscous	black glassy
RC-12C ОНС-ОУСНО 1.0 Н2	0 H ₂ N-(C)-NH ₂	1.0	үн ² ⊙-сно	4.0	brick- dust	granular	granular

Table 21
Elemental Analyses of Some of the RC-C
Series of Random Copolymers

Polymer	An	alyses,	%	
Number	an anno a transfer and an and an and an and an	H325		H400
	<u>C</u>	H	N	N
RC-3C	80.56	5.70	12.21	12.29
RC-4C	80.50	5.74	12.28	12.31
RC-5C	80.49	5.69	12.24	12.33
RC-6C	80.54	5.71	12.27	12.39
RC~7C	83.97	5.06	9.63	9.71
RC-8C	81.09	6.05	11.82	11.83
RC-9C	81.13	6.02	11.80	11.87
RC-10C	83.61	5.56	9.81	9.84
RC-11C	80.93	5.49	12.63	12.68

Table 22

Thermogravimetric Analyses of Some of the RC-C Series of H400
Random Copolymers in Nitrogen

Polymer Number	Atmos- phere	Per Cent Weight Loss at °C							
		400	500	600	700	800	900	1.000	1176
RC-3C-H400	nitrogen	0.0	0.0	2.6	4.8	8.1	13.5	14.7	17.5
RC-4C-H400	nitrogen	0.0	0.1	2.8	8.9	11.9	13.7	14.8	18.1
RC-5C-H400	nitrogen	0.0	0.0	3.0	9.4	12.3	14.3	15.7	18.9
RC-6C-H400	nitrogen	0.0	0.0	3.2	9.5	12.4	14.5	15.5	18.9
RC-7C-H400	ni.trogen	0.0	0.2	2.3	4.3	7.9	13.5	14.4	17.9
RC-8C-H400	nitrogen	0.0	0.3	1.9	4.1	7.7	13.3	14.7	17.4
RC-9C-H400	nitrogen	0.0	0.0	2.5	5.0	8.3	13.7	14.9	17.7
RC-10C-H400	nitrogen	0.0	0.0	2.9	9.1	12.1	14.0	15.2	18.2
RC-11C-H400	nitrogen	0.0	0.0	2.1	4.1	7.6	13.2	14.1	17.4

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